AD-A257 473



~ <u>4</u>

OFFICE OF NAVAL RESEARCH

Contract N00014-91-J-1910

R & T Code 4131025

Technical Report #47

Cathodic Reduction of Oxygen and Hydrogen Peroxide at Cobalt and Iron Crowned Phthalocyanines Adsorbed on Highly Oriented Pyrolytic Graphite Electrodes

By

N. Kobayashi, P. Janda and A.B.P. Lever*



in

Inorganic Chemistry

York University

Department of Chemistry, 4700 Keele St., North York

Ontario, Canada M3J 1P3

Reproduction in whole, or in part, is permitted for any purpose of the United States Government

*This document has been approved for public release and sale; its distribution is unlimited

*This statement should also appear in Item 10 of the Document Control Data-DD form 1473. Copies of the form available from cognizant contract administrator

92-28977

41204

92 11 05 028

REPORT DOCUMENTATION PAGE

Norm Approved
OMB No 0704-0138

1. AGENCY USE ONLY (Leave blank)	2. REPORT DATE		ND DATES COVERED	
	October 23, 1992	Technical ·	- June 1991 to July 1992	
TITLE AND SUBTITLE '			5. FUNDING NUMBERS	
Cathodic Reduction of Oxygen and Hydrogen Peroxide at			C - N00014-91-J-1910	
Cobalt and Iron Crowned	-			
Highly Oriented Pyrolyt	ic Graphite Electro	des	PR - 4131025	
AUTHOR(S)			i	
N, Kobayashi, P. Janda	and A.B.P. Lever			
PERFORMING ORGANIZATION NAME	S) AND ADDRESS(ES)		8. PERFORMING ORGANIZATION REPORT NUMBER	
A.B.P. Lever, Departme	nt of Chemistry			
York University, 4700		Parant #47		
North York, Ontario M3	J 1P3		Report #47	
Canada			Ì	
SPONSORING / MONITORING AGENCY	NAME(S) AND ADDRESS(ES)		10. SPONSORING MONITORING	
		AGENCY REPORT NUMBER		
Dr. Ronald A. De Marco				
Office of Naval Resear	ch, Chemistry Divis	ion	j	
800 N. Quincy St., Arl				
			74	
3. ABSTRACT (Maximum 200 words)				
The surface electrochemi (Fe ^{II} CRPc(-2) or Co ^{II} CRE graphite (HOPG) electrod The redox processes of s	c(-2)) adsorbed on he has been explored purface bound. Co ^{II} O	a highly orien under an argo: RPc(-2)/{Co ^I CR	ted pyrolytic n atmosphere. Pc(-2)] ⁻ ,	
${\tt [Fe^{III}CRPc(-2)]^+/Fe^{II}CRI}$	C(-2), and Fe ^{ll} CRPC r pH dependence rep	:(-2)/[Fe ⁺ CRPc(ported in the r	-2)] couples ange 1-13.	
Under an oxygen atmosphe	ere at the CoCRPC(-2)/HOPG elect	roue, two	
reduction processes, oxy	rgen to hydrogen per	oxide and nydr	a explored.	
to water were clearly de Surface bound Fe ^{II} CRPc(-	elineated, and their	ur electron re	duction of	
Surface bound FeCRPC(- oxygen in alkaline mediu	m. The Coll Cape / -2)/HOPG electro	de is very	
	ance and absence of	oxygen, but th	e Fe ^{II}	
oxygen in alkaline medic				
stable both in the prese	is much less stable	when electroc	atalytic oxygen	
stable both in the prese CRPc(-2)/HOPG electrode	is much less stable	when electroc	atalytic oxygen	
stable both in the prese CRPc(-2)/HOPG electrode	is much less stable	when electroc	atalytic oxygen	
stable both in the prese CRPc(-2)/HOPG electrode reduction takes place or	is much less stable	when electroc	atalytic oxygen	
stable both in the prese CRPc(-2)/HOPG electrode reduction takes place or	is much less stable	when electroc	atalytic oxygen	
stable both in the prese CRPc(-2)/HOPG electrode reduction takes place of SUBJECT TERMS	is much less stable its surface.	when electroc	15. NUMBER 25	
stable both in the prese CRPc(-2)/HOPG electrode reduction takes place or SUBJECT TERMS Phthalocyanine, Electrode	is much less stable its surface.	when electroc	atalytic oxygen	
stable both in the presence of the control of the c	is much less stable its surface.	when electroc	15. NUMBER 25	

TECHNICAL REPORT DISTRIBUTION LIST - GENERAL

Office of Naval Research (2)*
Chemistry Division, Code 1113
800 North Quincy Street
Arlington, Virginia 22217-5000

Dr. James S. Murday (1) Chemistry Division, Code 6100 Naval Research Laboratory Washington, D.C. 20375-5000

Dr. Robert Green, Director (1)
Chemistry Division, Code 385
Naval Air Weapons Center
Weapons Division
China Lake, CA 93555-6001

Dr. Elek Lindner (1)
Naval Command, Control and Ocean
Surveillance Center
RDT&E Division
San Diego, CA 92152-5000

Dr. Bernard E. Douda (1) Crane Division Naval Surface Warfare Center Crane, Indiana 47522-5000

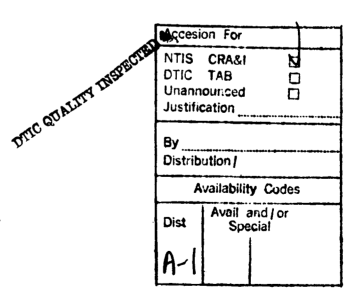
* Number of copies to forward

Dr. Richard W. Drisko (1)
Naval Civil Engineering
Laboratory
Code L52
Port Hueneme, CA 93043

Dr. Harold H. Singerman (1)
Naval Surface Warfare Center
Carderock Division Detachment
Annapolis, MD 21402-1198

Dr. Eugene C. Fischer (1)
Code 2840
Naval Surface Warfare Center
Carderock Division Detachment
Annapolis, MD 21402-1198

Defense Technical Information Center (2) Building 5, Cameron Station Alexandria, VA 22314



ABSTRACT DISTRIBUTION LIST

Professor Hector Abruña Department of Chemistry Cornell University Ithaca, NY 14853 Dr. Allen J. Bard
Department of Chemistry
University of Texas at Austin
Austin, TX 78712-1167

Professor Lesser Blum
Department of Physics
University of Puerto Rico
Rio Piedras, Puerto Rico 00931

Professor Daniel Buttry Department of Chemistry University of Wyoming Laramie, WY 82071

Professor Richard M. Crooks Department of Chemistry University of New Mexico Albuquerque, NM 87131 Professor Andrew Ewing
Department of Chemistry
152 Davey Laboratory
Pennsylvania State University
University Park, PA 16802

Professor Gregory Farrington
University of Pennsylvania
Department of Materials Science and Engineering
3231 Walnut Street
Philadelphia, Pennsylvania 19104

Professor W. R. Fawcett Department of Chemistry University of California, Davis Davis, CA 95616

Dr. John J. Fontanella Physics Department U.S. Naval Academy Annapolis, MD 21402-5026

Professor Harry Gray
California Institute of Technology
127-72
Pasadena, California 91125

Professor Joel Harris Department of Chemistry University of Utah Salt Lake City, UT 84112 Dr. Adam Heller
Department of Chemical Engineering
University of Texas at Austin
Austin TX 78712-1062

Professor Pat Hendra The University Southampton SO9 5NH England Professor Joseph Hupp Department of Chemistry Northwestern University Evanston, IL 60208

Professor Jiri Janata Department of Bioengineering University of Utah Salt Lake City, UT 84102 Professor A B. P. Level Department of Chemistry York University 4700 Keele Street North York, Ontario M3J 1P3

Professor Nathan S. Lewis Division of Chemistry and Chemical Engineering California Institute of Technology Pasadena, CA 91125 Dr. Bor Yann Liaw University of Hawaii at Manoa 2540 Maile Way, Spalding 253 Honolulu, HI 96822

Professor Rudolph Marcus Division of Chemistry and Chemical Engineering California Institute of Technology Pasadena, CA 91125 Professor Charles Martin Department of Chemistry Colorado State University Ft. Collins, CO 80523 Dr. Donald Maricle International Fuel Cells P. O. Box 739 195 Governors Highway South Windsor, CT 06074 Dr. Melvin H. Miles Energetic Materials Branch, Code 3853 Chemistry Division, Research Department Naval Weapons Center China Lake CA 93555

Professor Royce W. Murray Department of Chemistry University of North Carolina at Chapel Hill Chapel Hill, NC 27514 Dr. David J. Nagel Naval Research Laboratory Code 4600 4555 Overlook Avenue, S.W. Washington, D.C. 20375-5000

Dr. Michael R. Philpott IBM Research Division Almaden Research Center 650 Harry Road San Jose, CA 95120-6099 Professor B. S. Pons Department of Chemistry University of Utah Salt Lake City, UT 84112

Dr. Mark A. Ratner
Department of Chemistry
Northwestern University
Evanston, IL 60208

Dr. Debra Rolison Code 6170 Naval Research Laboratory Washington, DC 20375-5000

Dr. Michael J. Sailor Department of Chemistry University of California, San Diego 9500 Gilman Drive La Jolla CA 92093-0506 Professor Jack Simons
Department of Chemistry
University of Utah
Salt Lake City, UT 84112

Professor John L. Stickney University of Georgia Department of Chemistry Cedar Street Athens, GA 30602 Professor Eric M. Stuve
Dept. of Chemical Engineering, BF-10
University of Washington
Seattle, Washington 98195

Dr. Stanislaw J. Szpak Code 574 Naval Ocean Systems Center San Diego, CA 92152-5000 Dr. E. Jennings Taylor Physical Sciences, Inc. 20 New England Business Center Andover MA 01810

Dr. Petr Vanysek Department of Chemistry Northern Illinois University DeKalb, IL 60115 Professor Michael Weaver Department of Chemistry Purdue University West Lafayette, IN 49707

Professor Henry S. White
Department of Chemical Engineering
and Materials Science
421 Washington Avenue, SE
University of Minnesota
Minneapolis, MN 55455

Professor R. Mark Wightman
Department of Chemistry
CB #3290, Venable Hall
The University of North Carolina
Chapel Hill NC 27599-3290

Professor George Wilson Department of Chemistry University of Kansas Lawrence, KS 66045 Professor Mark S. Wrighton
Department of Chemistry
Massachusetts Institute of Technology
Cambridge, MA 02139

Professor Ernest Yeager
Case Center for Electrochemical Sciences
Case Western Reserve University
Cleveland, OH 44106

Cathodic Reduction of Oxygen and Hydrogen Peroxide at Cobalt and Iron Crowned Phthalocyanines
Adsorbed on Highly Oriented Pyrolytic Graphite Electrodes

Nagao Kobayashi*¹, Pavel Janda,*², and A.B.P. Lever*
Department of Chemistry, York University, 4700 Keele St.
North York (Toronto), Ontario, Canada M3J 1P3

The surface electrochemistry of iron and cobalt crowned phthalocyanine (Fe^{II}CRPc(-2) or Co^{II}CRPc(-2)) adsorbed on a highly oriented pyrolytic graphite (HOPG) electrode has been explored under an argon atmosphere. The redox processes of surface bound, Co^{II}CRPc(-2)/[Co^ICRPc(-2)]-, [Fe^{III}CRPc(-2)]+/
Fe^{II}CRPc(-2), and Fe^{II}CRPc(-2)/[Fe^ICRPc(-2)]- couples were identified and their pH dependence reported in the range 1-13. Under an oxygen atmosphere at the Co^{II}CRPc(-2)/HOPG electrode, two reduction processes, oxygen to hydrogen peroxide and hydrogen peroxide to water were clearly delineated, and their pH dependences explored. Surface bound Fe^{II}CRPc(-2) catalyses the four electron reduction of oxygen in alkaline medium. The Co^{II}CRPc(-2)/HOPG electrode is very stable both in the presence and absence of oxygen, but the Fe^{II}CRPc(-2)/HOPG electrode is much less stable when electrocatalytic oxygen reduction takes place on its surface.

Introduction

There has been significant progress in our understanding of the electrocatalytic reduction of oxygen in the presence of organometallic catalysts³⁻⁶ In many cases, oxygen reduction has been carried out at catalyst-adsorbed or -modified electrodes. When monomeric cobalt porphyrins and phthalocyanines are used as catalysts, oxygen reduction usually proceeds only to hydrogen peroxide $(H_2O_2)^{3,4}$, however the four-electron reduction of oxygen to water catalysed by a monomeric cobalt phthalocyanine⁷ and by a cyano bridged cobalt phthalocyanine⁸ has recently been reported. Some cobalt derivatives of cofacial diporphyrin are known to catalyze the reduction of oxygen to water⁵. Moreover, iron porphyrins and phthalocyanines will catalyse the four-electron reduction of oxygen, in alkaline medium. However, the factors responsible for activating a four-electron rather than a two-electron pathway, remain obscure. Oxygen reduction characteristics are influenced by various extrinsic factors such as, for example, differences in adsorption procedure and of surface coverage by catalyst, and even by changes of ionic strength of the experimental solution^{6k}. Many catalysts are degraded by contact with hydrogen peroxide such that successive potential scans of a given modified electrode surface are not fully reproducible, necessitating a large number of studies with fresh surfaces. It is difficult to ensure that all such fresh surfaces are identical further complicating the study.

Recently, we reported the synthesis and properties of phthalocyanines with four 15-crown-5 ether voids at the 3,4 position⁹, $MCrPc(-2)^{10}$ (M = Co, Fe, Cu, H₂). In the course of purification, we noted that these phthalocyanines adsorbed onto silica-gel and powdered charcoal very strongly. Also, they exhibited ideal spectroscopic (UV, MCD, ESR, NMR, and Emission etc.) behavior. The cobalt and iron derivatives (Figure 1) have therefore been studied as electrocatalysts for oxygen reduction, and are shown here to have many desirable features and in particular they (Fe,Co) can be used for preparation of stable electrodes with modified surfaces whose electrochemical properties are sufficiently reproducible for multiple scan voltammetry. It is not necessary to constantly renew the surface, at least under argon. The iron derivative, under oxygen, is less stable. The cobalt species is more stable but shows some deactivation over time when deposited on the RDE while

electrocatalytic oxygen reduction takes place.

During the progress of our studies, there has evidently been great interest in the properties of both crowned phthalocyanines ¹¹ and crowned porphyrins ¹² but the properties of these with respect to oxygen electrocatalysis have not been explored.

Co^{II}CRPc(-2) adsorbed onto highly oriented pyrolytic graphite electrodes (HOPG)¹³ is shown to catalyse the commonly observed two-electron reduction of oxygen to hydrogen peroxide. This is followed at a significantly more negative potential by the further reduction of hydrogen peroxide to water. While such a double step reduction has been observed previously,³g,⁶g,¹³⁻¹⁶, the very clean separation observed here between these two processes affords us the rare opportunity of a more detailed study of the pH dependence.

Electrocatalytic data for Fe^{II}CRPc(-2) adsorbed onto a HOPG electrode are also described. As with other iron phthalocyanines, Fe^{II}CRPc(-2) catalyzes the four-electron reduction to water at alkaline pH. Interestingly, the pH dependence for oxygen reduction at the Fe^{II}CRPc(-2) adsorbed electrode is quite similar to that of hydrogen peroxide reduction at the Co^{II}CRPc(-2)/HOPG electrode. However the data are not of such high quality as with Co^{II}CRPc(-2)/HOPG due to decomposition of the Fe^{II}CRPc(-2).

Experimental Section

Materials. Argon gas (Linde) was purified by passage through heated copper filings, anhydrous CaSO₄ (Drieri*e), molecular sieves (Bolf, Type 3A), and glass wool. Oxygen gas (Linde) was purified by passage through anhydrous CaSO₄, NaOH pellets (AnalaR analytical grade), anhydrous CaSO₄, molecular sieves and glass wool.

The cobalt crown phthalocyanine, Co^{II}CRPc(-2) was prepared according to a literature procedure. ^{9a,9b}. The iron crown phthalocyanine, Fe^{II}CRPc(-2) was prepared and purified in a similar fashion, ^{8b} from 2,3-(3',4'-dicyanobenzo)-1,4,7,10,13-pentaoxacyclopentadeca-2-ene and FeCl₂ in refluxing ethylene glycol. <u>Anal.</u> Calcd for C₆₄H₇₂FeN₈O₂₀: C,57.83; H,5.46; N,8.43. Found: C,57.59; H,5.40; N,8.38.

Electrochemical measurements: These were undertaken using equipment (Pine 4-electrode potentiostat, Princeton Applied Research Corporation, PARC potentiostat) and techniques as previously described. The electrode was mounted in Teflon to expose a circular area of 0.440 cm². Prior to adsorption of the catalyst, the HOPG (Union Carbide) surface was renewed using adhesive Scotch Tape. A freshly exposed HOPG surface was modified by dipping and rotating it in 0.1mM Fe^{II}CRPc(-2) or Co^{II}CRPc(-2) in DCB/ethanol (19:1 v/v) solution for about 5 - 10 min. The electrode was then washed with ethanol and doubly distilled water. Current-potential (i - E) curves were recorded in aqueous 0.1M phosphate buffer (prepared from 0.1M solution of Na₂HPO₄ and KH₂PO₄), the pH was adjusted by addition of small amount of KOH and H₃PO₄(AnalaR, BHD chemicals) as required. Measurements at pH 1 - 2 were carried out in dilute H₂SO₄ (Suprapur, Merck).

Results and Discussion

- i) Studies under an Argon atmosphere, in the absence of Oxygen.
- a) Voltammetric Response of Co^{II}CRPc(-2)/HOPG.

Figure 2A shows cyclic voltammetric (CV) potentiodynamic waves obtained at bare HOPG and Co^{II}CRPc(-2)/HOPG electrodes, with data collected in Table 1. An anodic and a cathodic peak of practically equal area appeared in curve b, with peak currents (i_p) directly proportional to scan rate, typical of a Faradaic process involving an adsorbed species. Assuming that the surface redox couple obeys the Nernst equation, the number of electrons participating per adsorbed molecule also estimated from the slope of the i_p versus scan rate plot is 0.972 ± 0.002^{18} , i.e. a one-electron redox process. The charge under this peak is then $4.1 \,\mu$ C and corresponds to a surface concentration of $4.2 \times 10^{-11} \text{mol/cm}^2$. From the Corey-Pauling-Koltun molecular model, the length of one side of Co^{II}CRPc(-2) is about 20 Å. Using the above surface concentration and assuming that Co^{II}CRPc(-2) lies flat on the electrode, the calculated coverage corresponds to almost exactly one monolayer. This limiting coverage of adsorbed Co^{II}CRPc(-2) was independent of the concentration of Co^{II}CRPc(-2) in the DCB used during adsorption. The Co^{II}CRPc(-2)/HOPG electrode was stable after several hundreds cycles in the potential range, shown in Figure 2, in the

whole pH range and was not affected by rotation of the electrode.

The pH dependence of the redox potential calculated as a mean value of anodic and cathodic peak potential is shown in Figure 3B, and Tables 1,2. High regression coefficients provide some confidence for the accuracies of the slopes and intercepts. In the range pH 5 - 1.6, the potential shifts to a more negative direction with increasing pH with slope of about -63 mV/pH, suggesting participation of one proton. Above pH 5, it becomes essentially pH-independent. The potential values, and their dependence on pH, are fully comparable to earlier studies with Co^{II}TNPc(-2) (TNPc(-2) = tetraneopentoxyphthalocyanine)¹³ and Co^{II}TsPc(-2) (TsPc(-2) = tetrasulfonylphthalocyanine)^{3h,19} and the wave in question undoubtedly arises from the Co^{II}/Co^I couple.

During our variable pH study, some experiments revealed a wave at more positive potentials (as noted in Table 1). This was generally poorly developed. It is almost certainly the [Co^{III}CRPc(-2)] +/ Co^{II}CRPc(-2) wave but its identification was not pursued.

In our studies with cobalt tetrasulfonated phthalocyanine, ¹⁹ Co^{II}TsPc(-2)/HOPG, a second reduction process was observed occurring negative of the Co^{II}TsPc(-2)/[Co^ITsPc(-2)]⁻ couple, namely the [Co^ITsPc(-2)]⁻/[Co^ITsPc(-3)]²⁻ process. This is observed relatively close to the former process in acid medium, but its different pH dependence causes it to be separated therefrom in basic medium. The pH dependence data for the latter process are reproduced in Figure 3C since they are relevant to the discussion below. In the Co^{II}CRPc(-2) case, this second reduction process was detected in strong acid medium (see Table 1). However Co^{II}CRPc(-2) catalyses proton (water) reduction and the solvent cut-off through this process obscures this second reduction step for all solutions except those with pH near 1.

For completion note that a Co^{II}CRPc(-2)/HOPG electrode studied in DCB/TBAP solution showed a cyclic voltammogram, using a RDE electrode, with surface-bound couples appearing at -0.53 V (vs SCE) Co^{II}CRPc(-2)/[Co^ICRPc(-2)]⁻ and at +0.32 and +0.46 V attributable to the couples²⁰ [Co^{II}CRPc(-1)] +/Co^{II}CRPc(-2) and [Co^{III}CRPc(-1)]² +/[Co^{II}CRPc(-1)] + respectively.

b) Voltammetric Response of Fe^{II}CRPc(-2)/HOPG.

In oxygen-free solution, a Fe^{II}CRPc(-2)/HOPG electrode shows two redox couples (Figure 2B) in the potential range from ca. +0.2 V to -0.4 V, with pH-dependence as shown in Figure 4A. From comparison with literature data on pH dependence, 6e,19,21 the more positive potential couple is assigned to [Fe^{III}CRPc(-2)] + /Fe^{II}CRPc(-2) and the less, to Fe^{II}CRPc(-2)/[Fe^ICRPc(-2)]⁻. Although the [Fe^{III}CRPc(-2)] + /Fe^{II}CRPc(-2) couple looks sharper than that of Co^{II}CRPc(-2)/[Co^ICRPc(-2)]⁻ in Figure 2A, the actual surface concentration of Fe^{II}CRPc(-2) (4.8 x $^{10-11}$ mole/cm²) estimated from the charge involved in the process (4.0 μ C/cm²) is approximately the same as that of Co^{II}CRPc(-2), i.e. a monolayer. Regression data are collected in Table 2.

At pH above 7, it was not possible to observe the Fe^{II}CRPc(-2)/[Fe^ICRPc(-2)]⁻ wave due to solvent limit (proton reduction). It is clearly observed ¹⁹ with Fe^{II}TsPc(-2) which must evidently be less effective in catalysing proton reduction.

c) Studies using Hydrogen Peroxide and Co^{II}CRPc(-2)/HOPG.

Cyclic voltammetric studies were briefly undertaken, of the reduction of oxygen-free, argon saturated, H_2O_2 at a bare HOPG electrode, and at a $Co^{II}CRPc(-2)/HOPG$ electrode, for comparison with the oxygen reduction data below. The bare electrode shows rising reduction current at -1.0 V, at pH 3, while the $Co^{II}CRPc/HOPG$ electrode shows a reduction peak at -0.88 V corresponding almost exactly with the second oxygen reduction process described below.

- ii) Studies in Oxygen Saturated Buffer Solutions.
- a) Voltammetric Response of Co^{II}CRPc(-2)/HOPG.

Figure 5A displays the CV curves for the reduction of O_2 . Two cathodic peaks were recognized before the current due to H_2 evolution obscured further processes. The first is due to the reduction of oxygen to H_2O_2 , and the second to the reduction of the H_2O_2 so produced. The second peak corresponds with that observed when hydrogen peroxide is studied alone, (see immediately above) and increases in current if hydrogen peroxide is added to the oxygenated solution.

and second cathodic peak potential permitting clear definition of the pH dependence of both peak potentials (vide infra).

Figure 5B shows the i - E responses for the first oxygen reduction wave obtained at a Co^{II}CRPc(-2)/HOPG rotating disk electrode (RDE) at several pH values. Multiple potential scans of the same surface give good reproducible data. The onset of oxygen reduction and the limiting current depend upon the solution pH. However, this last observation is due to the difference of solubility of oxygen: greater in basic than in acidic solution. The number of electrons, n, involved in the reaction was calculated from the Levich equation: 13,18,22

$$i_l = 0.62 \text{ nFA}\omega^{1/2} C_{ox} D^{2/3} \nu^{-1/6}$$
 (1)

where C_{OX} is the bulk oxygen concentration, D_O is its diffusion coefficient, ν is the kinematic viscosity and the other symbols have their usual electrochemical meanings. A value of $C_{OX}D^{2/3}\nu^{-1/6}$ = 1.77 x 10⁻⁹ was derived via calibration with a platinum electrode (assuming a well behaved 4-electron reduction process at Pt),²³ and led to an average experimental value of 1.99 ± 0.03 electrons, confirming that oxygen is reduced to H_2O_2 .

RDE data show (Figure 5C) linear dependence of i_1 on the square root of the rotation rate indicating a fast diffusion driven reaction. Note the rising current (Figure 5B), beyond the plateau, at more negative potentials corresponding to the ensuing reduction of H_2O_2 , and especially noticeable at lower pH. The data were analyzed by plotting them as i^{-1} vs $\omega^{-1/2}$ (Figure 4E) (Koutecky-Levich plots 13 ,22). The line is almost perfectly straight, indicating first-order dependence of the overall reaction on oxygen concentration. The intercept is indistinguishable from zero within experimental error indicating a very fast reaction rate. Figure 5E shows a Tafel type plot from RDE data, for the first reduction process, at several representative pHs. The slope of the plot changes slightly in the pH range of 1.6 to 5 from -116 to -130 mV/decade, but above pH 5 this value remains constant (-130 mV/decade). This may be indicative of an irreversible (non-Nernstian) process possibly with a one-electron rate determining step. Curve A in Figure 3 represents the pH dependence of the half-wave potential, $E_{1/2}$, for oxygen reduction at $Co^{II}CRPc(-2)/HOPG$. The $E_{1/2}$ for oxygen reduction is always more positive than the Co^{II}/Co^{I}

 $Co^{II}CRPc(-2)/HOPG$. The $E_{1/2}$ for oxygen reduction is always more positive than the Co^{II}/Co^{I} process (Figure 3, curve B), but with a much shallower pH dependence, -33 mV/pH, in the pH regime 1-4.

The pH dependence for reduction of hydrogen peroxide to water at a Co^{II}CRPc(-2)/HOPG electrode is shown in Figure 3D. It differs substantially from that of oxygen reduction at Co^{II}CRPc(-2)/HOPG, shown in Figure 3A. There are three regions of differing slope showing that the number of protons involved in the process varies as the mechanism varies from low to medium to high pH. At low pH, the slope is -110 mV/pH. Near neutral pH, the data are more scattered but the slope is evidently zero or small. Unfortunately it was difficult to collect data for this process at high pH since the process occurs very close to the reduction of water.

b) Mechanism of Oxygen and Hydrogen Peroxide Reduction by Co^{II}CRPc(-2)

It is tempting to assume that the active catalyst for oxygen reduction is the [Co^ICRPc(-2)]⁻ species since the pH dependence of oxygen reduction parallels that of the Co^{II}Pc(-2)/[Co^IPc(-2)]⁻ couple, at slightly more positive potentials. Indeed with a simplified assumption of reversibility, and ideal behavior of the phthalocyanine catalyst, one may calculate a rate constant for oxygen reduction,²⁴ using:

$$k = (D_0/\Gamma_{CoCRPc}\delta) \exp[(F/RT)(E_{1/2}-E^0)]$$
 (2)

where δ is the thickness of the diffusion layer at the RDE (and $\delta=4.98D_0^{1/3}\nu^{1/6}\omega^{1/2})^{24c}$, Γ_{CoCRPc} is the surface concentration of the catalyzer, $E_{1/2}$ is the half-wave potential for oxygen reduction, and E^0 is the redox potential for the $Co^{II}CRPc(-2)/[Co^ICRPc(-2)]^-$ redox couple. The values (T = 20° C) $D_0 = 1.7 \times 10^{-5}$ cm²/s, $\Gamma = 4.2 \times 10^{-11}$ mol/cm², rotation rate $\omega = 42$ rads/s, kinematic viscosity $\nu = 10^{-2}$ cm²/s and the value of $E_{1/2} - E^0 = ca$ 300 mV, from the data, yield k = ca 10^{10} M⁻¹s⁻¹ in basic medium. However this does not prove the catalytic participation of $Co^IPc(-2)$. Recently we have discussed the electrocatalytic reduction of oxygen by perchlorinated cobalt phthalocyanine ($CoCl_{16}Pc(-2)$). In this case, oxygen reduction parallels the $Co^{II}Pc(-2)/[Co^IPc(-2)]^-$ couple in acid medium, but not in alkaline medium. In strong alkali, oxygen reduction occurs at some 400 - 600 mV more positive a potential than the $Co^{II}Pc(-2)/[Co^IPc(-2)]^-$

couple. If the [Co^ICl₁₆Pc(-2)]⁻ were the active species, oxygen reduction would have to occur at an unrealistically high rate (following the above analysis). Therefore we proposed that the active species was Co^{II}Cl₁₆Pc(-2). It is reasonable to suppose that Co^{II}CRPc(-2) is the active catalyst here and that an intermediate oxygen adduct is involved: ^{13,25}

$$Co^{II}Pc(-2) + O_2 < = = = = > (O_2^-)Co^{III}Pc(-2)$$
 (3)

wherein in (3), the most simple oxygen adduct is portrayed, based on cobalt(III) dioxygen chemistry. A binuclear peroxo-bridged species might also be invoked, though, in the surface environment, a mononuclear species is perhaps more readily conceived. The argument remains substantially equivalent if a mononuclear or a binuclear oxygen adduct is assumed. Equilibrium (3) is strongly driven to the left and therefore oxygen reduction is not generally observed at, or close to, the [Co^{III}Pc(-2)] +/Co^{II}Pc(-2) couple potential, nor are oxygen adducts seen when Co^{II}Pc(-2) solutions are mixed with oxygen. However oxygen adducts have been observed at low temperatures, with CoPc, proving that equilibria such as (3) do occur. 27

The data are consistent with the supposition that reduction of the oxygen adduct in (3) occurs at the potential corresponding experimentally with the observed catalytic oxygen reduction process. Removal of the oxygen adduct via this reduction process then drives the equilibrium (3) to the right. The pH dependence follows the Co^{II}/Co^I couple in the acid regime for two possible reasons, i) protonation of the peroxo bridge likely occurs, and ii) the [Co^ICRPc(-2)]⁻ species is air (oxygen) sensitive so that it would also catalyse oxygen reduction at a potential close to the Co^{II}/Co^I process as [Co^ICRPc(-2)]⁻ begins to build up. The involvement of an analogous Co^{II} porphyrin as the catalytically active species has also recently been suggested by Anson and co-workers²⁸.

The reduction of hydrogen peroxide occurs at a potential somewhat negative of that for the next phthalocyanine reduction couple, $[Co^{I}Pc(-2)]^{-/}[Co^{I}Pc(-3)]^{2-}$ which was tracked as a function of pH in a parallel study of CoTsPc¹⁹ (Figure 3C). Thus the species $[Co^{I}CRPc(-3)]^{2-}$ will be the effective catalyst in peroxide reduction, probably via a complex similar to that described in (3).

This last couple is more accurately portrayed as HCo^IPc(-2)/H₂Co^IPc(-3), in acid medium, where protonation (probably at a bridgehead aza link, but conceivably also at the metal center) has

occurred and where the number of protons is dictated by consideration of the pH dependences of the Co^{III}/Co^{II}, Co^{II}/Co^I and Co^IPc(-2)/Co^IPc(-3) couples shown in Figure 3. Assuming Co^{II}Pc is not protonated, then the degree of protonation indicated above follows. In basic medium, [Co^ICRPc(-2)]⁻ will then be unprotonated, while [HCo^ICRPc(-3)]⁻ will retain one proton. Thus the reduction of hydrogen peroxide will be catalysed by these protonated [Co^IPc(-3)]²⁻ species. Presentation of a detailed mechanism awaits further study. However note that mechanisms have been invoked to include coordination of HO₂⁻ to a reduced cobalt phthalocyanine, followed by bond breaking and generation of a hydroxide radical attached to cobalt, which is then immediately reduced to coordinated hydroxide anion.³ⁱ

b) Voltammetric Response of Fe^{II}CRPc(-2)/HOPG.

At low pH, oxygen reduction starts (Figure 4B) slightly negative of the potential of the [Fe^{III}CRPc(-2)] + /Fe^{II}CRPc(-2) reduction couple (also see Figure 6A, curves a and b), strongly suggesting the participation of Fe^{II}CRPc(-2) in the catalytic reduction process. The pH dependence for oxygen reduction (Figure 4B) roughly tracks that of the [Fe^{III}CRPc(-2)] + /Fe^{II}CRPc(-2) couple (Figure 5A). The separation between the [Fe^{III}CRPc(-2)] + /Fe^{II}CRPc(-2) and the oxygen reduction process is proportional to the rate of oxygen reduction at the modified surface. Thus consideration of Figure 4 shows that the rate increases between about pH 3 and 7 and then is approximately independent of pH.

The i - E responses recorded at the Fe^{II}CRPc(-2)/HOPG RDE in pure oxygen saturated solution (pH 9.2) (Figure 6B) were analysed by the conventional Koutecky-Levich²² approach. The data are similar to those observed with FeTsPc.⁶ⁱ A Levich plot (not shown) of i_1 vs $\omega^{1/2}$ from these curves (Figure 6C) is linear with regression coefficient 0.999. An average value of n of 3.83 was obtained from the Levich equation (Eqn. (1) above). The currents at the maxima then correspond to pure diffusion control for a four-electron reduction. The pH dependence for oxygen reduction (Figure 4B) is remarkably similar to that for H_2O_2 reduction catalysed by CoCRPc (Figure 3D) suggesting that reduction may follow a similar mechanism at least insofar as proton dependence is concerned.

Thus, as for several other iron phthalocyanines^{4,6i,6j}, Fe^{II}CRPc(-2) catalyzes the overall four-electron reduction in alkaline solution (in acidic solution, the catalytic current is smaller both in CV and RDE i - E diagrams).

Concluding Remarks

The crown phthalocyanine iron and cobalt species presents reproducible and well behaved oxygen and hydrogen peroxide reduction data. In this preliminary report, we present an overview of these properties.

The excellent pH dependence data reported here, and the high stability of the surface, suggest that the CoCRPc system, in particular (the FeCRPc surface is less stable) could provide much more detailed mechanistic information concerning the catalytic processes involving the reduction of oxygen, hydrogen peroxide and indeed protons. The pH studies reported here may only be interpreted in rather simple terms in the absence of a more definitive study of the various steps which participate in what is likely a complex mechanism. However the study would be worthy of expansion to other crown phthalocyanine species such as those with manganese, rhodium and iridium. Studies involving the oxygen reduction of cobalt and iron crown phthalocyanine species as a function of the cation contained within the crown residue, are in progress.

Acknowledgements: We are indebted to the Natural Sciences and Engineering Research Council (Ottawa) and the Office of Naval Research (Washington) for financial support. We are also pleased to acknowledge the technical assistance of Penny Seymour. We are indebted to Union Carbide (Parma) for the supply of HOPG.

Table 1 Redox Potentials for CoPc, Oxygen and Hydrogen Peroxide Redox Processes, vs SCE.a,b

	Co ^{II} Pc(-2)	[Co ^I Pc(-2)] ⁻ /		<u> </u>	Ч-О-
				O ₂	H ₂ O ₂
	\[Co ¹ Pc(-2)] ⁻	$/[Co^{I}Pc(-3)]^{2}$		$/H_2O_2$	/H ₂ O
pН	E _{1/2}	E _{1/2}	pН	Ep	Ep
	CRPc	TsPc		CRPc	CRPc
1.0	-0.34		1.0	-0.29	
1.6	-0.38	-0.48	1.6	-0.30	-0.80
2.1	-0.42	-0.51	2.0	-0.29	-0.84
3.0	-0.46	-0.57	2.1	-0.32	-0.84
4.0	-0.54	-0.64	3.0	-0.35	-0.94 ~ _{**}
5.0	-0.59	***	4.0	-0.38	-1.06
5.7	-0.62	***	5.0	-0.37	-1.06
6.0	-0.60		5.7	-0.35	-1.08
7.0	-0.60	-1.04	6.0	-0.36	-1.15
8.1	-0.60	-1.10	7.0	-0.37	-1.15
9.0	-0.60	-1.15	8.1	-0.32	-1.34
10.0	-0.60	-1.21	9.0	-0.32	-1.39
11.0	-0.59	-1.26	10.0	-0.33	
12.0	-0.62	-1.32	11.0	-0.33	-1.40
13.0	-0.62	-1.37	13.0	-0.37	

a) All data are from cyclic voltammetry. Averages of the anodic and cathodic waves of the reversible metal centered couples are quoted. The irreversible cathodic peak potential is quoted for the oxygen reduction processes. b) Additional data: A second Co^{II}CRPc(-2) reduction potential was observed at -0.47 V for pH = 1. The oxidation process forming [Co^{III}CRPc(-2)] + was observed at 0.67, 0.52 and 0.28 at pH 1,7 and 10 respectively.

Table 2 Regression Data for pH Dependence (Data in volts, vs SCE)

Redox Process	pH Range	Slope ^a	Intercept ^a	R	# ²
Co ^{II} CRPc(-2)	1.6 - 5	-0.063(.002)	-0.28(.008)	0.997	6
/[Co ^I CRPc(-2)]	5 - 13	0p			
$[Co^{I}TsPc(-2)]^{-}$	1.25 - 4.5	-0.054(.006)	-0.41(.015)	0.988	4
$/[Co^{I}TsPc(-3)]^{2}$	7.4 - 13	-0.056(.002)	-0.64(.01)	0.997	6
$O_2> H_2 O_2$	1.6 - 4	-0.033(.005)	-0.25(.012)	0.950	6
	5 - 13	0p			
$H_2O_2 > H_2O$	1.6 - 7	-0.110(.004)	-0.62(.008)	0.998	5
[Fe ^{III} CRPc(-2)] +	0.9 - 5	0	+0.18		
/Fe ^{II} CRPc(-2)	7 - 13	-0.054(.001)	+0.45(.007)	0.999	5
Fe ^{II} CRPc(-2)	1-5	-0.089(.005) ^c	-0.12(.016)	0.994	6
/[Fe ^I CRPc(-2)]					
Oxygen reduction	1 - 4	-0.049(.007)	+0.17(.017)	0.952	7
	8 - 13	-0.034(.005)	+ 0.12(.020)	0.953	7

a) Least squares analysis using number of points as indicated and with regression coefficient as indicated. b) Assumed zero slope; in fact a least squares analysis would yield a very small negative slope. c) Omission of the point at pH = 7, provides a better correlation with a slope of -0.069 V/pH.

Table 3 Redox Potentials for Fe^{II}CRPc Surface Potentials under Argon and Under Oxygen. vs SCE

	[Fe ^{III} CRPc(-2)] +		Fe ^{II} CRPc(-2)		Oxygen	
	/Fe ^{II} CRPc(-2)	/[Fe ^I CRPc(-2)]	Reduction			
pН	E _{1/2}	E _{1/2}	pН	E _p		
0.9	0.16	-0.21	1.0	0.13		
1.3	0.21	-0.23	1.3	0.08		
2.2	0.17	-0.34	1.3	0.13		
3.5	0.15	-0.43	2.0	0.07		
4.0	0.19	-0.46	2.7	0.03		
4.6	0.15	-0.54	3.3	0.02	-4	
3.5	0.00		3.5	0.00		
7.0	0.08		4.0	0.00		
4.6	0.00		4.6	0.00		
5.5	-0.03		5.5	-0.30		
7.0	-0.06		7.0	-0.06		
8.9	-0.03	·	7.9	-0.07		
10.3	-0.11		8.3	-0.13		
12.0	-0.20		8.3	-0.13		
13.0	-0.24		8.9	-0.20		
9.3	-0.22		9.3	-0.22		
10.3	-0.22		10.3	-0.22		
11.1	-0.26		11.1	-0.26		
12.0	-0.28		12.0	-0.28		
			13.0	-0.32		

See footnote a) in Table 1.

Bibliography

- 1. Visiting professor from the Pharmaceutical Institute, Tohoku University, Sendai 980, Japan.
- 2. Visiting scholar from the J. Heyrovsky Institute of Physical Chemistry and Electrochemistry, Czechoslovak Academy of Sciences, Praha, Czechoslovakia.
- 3. (a) Jahnke, H.; Schonborn, M.; Zimmerman, G. Topics in Current Chemistry 1976, 61, 133. (b)

 Tarasevich, M.R.; Radyushkina, K.A. Russ. Chem. Chem. Eng. Trans. 1980, 49, 1498. (c) Van den Brink, F.; Barendrecht, E.; Visscher, W. Rec. J. Roy. Neth. Chem. 1980, 99, 253. (d) Van Veen, J.A.R.; Van Baar, J.F. Rev. Inorg. Chem. 1982, 4, 293. (e) Schiffrin, D.J.

 Electrochemistry 1983, 8, 126. (f) Yeager, B. Electrochim. Acta 1984, 29, 1527; J. Mol.

 Catal. 1986, 38, 5. (g) Elzing, A.; Van Der Putten, A.; Visscher, W.; Barendrecht, E. J.

 Electroanal. Chem. 1986, 200, 313. (h) Van Der Putten, A.; Elzing, A.; Visscher, W.;

 Barendrecht, E. J. Electroanal. Chem. 1987, 221, 95. (i) Van den Brink, F.; Visscher, W.;

 Barendrecht, E. J. Electroanal. Chem. 1983, 157, 305.
- 4. Kobayashi, N.; Nishiyama, Y. J. Phys. Chem. 1985, 89, 1167 and many refs. therein.
- (a) Collman, J.P.; Marrocco, M.; Denisevich, P.; Koval, C.; Anson, F.C. <u>J. Electranal. Chem.</u>
 1979, 101, 117; <u>J. Am. Chem. Soc.</u> 1980, 102, 6027. (b) Chang, C.K.; Abdalmuhdi, I. <u>J. Am.</u>
 <u>Chem. Soc.</u> 1984, 106, 2725. (c) Durand, R.R.: Bencosme, C.S.; Collman, J.P.; Anson, F.C. <u>J. Am. Chem. Soc.</u> 1983, 105, 2710.
- (a) Kobayashi, N.; Fujihira, M.; Sunakawa, K.; Osa, T. J. Electroanal. Chem. 1979, 103, 427. (b)
 Kuwana, T.; Bettelheim, A. Anal. Chem. 1979, 51, 2257. (c) Kuwana, T.; Chan, R.J.;
 Bettelheim, A. J. Electroanal. Chem. 1980, 110, 93. (d) Kuwana, T.; Forshey, P.A. Inorg.
 Chem. 1983, 22, 699. (e) Kuwana, T.; Forshey, P.A.; Kobayashi, N.; Osa, T. Adv. Chem. Ser.
 1982, No. 201, 601. (f) Kobayashi, N.; Nishiyama, Y. J. Electroanal. Chem. 1984, 181, 107. (g)
 Shigehara, K.; Anson, F.C. J. Phys. Chem. 1982, 86, 2776. (h) Wan, G-X.; Shigehara, K.;
 Tsuchida, E.; Anson, F.C. J. Electroanal. Chem. 1984, 179, 239. (i) Zagal, J.; Bindra, P.;
 Yeager, E. J. Electrochem. Soc. 1980, 12, 1506. (j) Kozawa, A.; Zilionis, V.E.; Brodd, R.J.

- ibid. 1970, 117, 1470; 1971, 118, 170. (k) Van den Brink, F.; Visscher, W.; Barendrecht, E. J. Electroanal. Chem. 1984, 172, 301; 1984, 175, 279. Zagal, J.; Sen, R. K.; Yeager, E. J. Electroanal. Chem. 1977, 83, 207.
- 7. Kobayashi, N.; Sudo, K.; Osa, T. <u>Bull. Chem. Soc. Jpn</u> 1990, <u>63</u>, 571.
- 8. Ikeda, O.; Itoh, S.; Yoneyama, H. Bull. Chem. Soc. Jpn. 1988, 61, 1428.
- (a) Kobayashi, N.; Nishiyama, Y. <u>J. Chem. Soc., Chem. Commun.</u> 1986, 1462. (b) Kobayashi, N.;
 Lever, A.B.P. <u>J. Am. Chem. Soc.</u> 1987, 109, 7433. (c) Gasyna, Z.; Kobayashi, N.; Stillman, M.
 J., <u>J. Chem. Soc., Dalton Trans</u> 1989, 2397.
- 10. Abbreviations: FeCRPc(-2), tetra(15-crown-5)ed phthalocyaninatoiron; CoCRPc(-2), tetra(15-crown-5)ed phthalocyaninatocobalt; HOPG, highly oriented pyrrolytic graphite; CV, cyclovoltammetric; RDE, rotating disk electrode; SCE, saturated calomel electrode; DCB, dichlorobenzene; i, current;
- Gurek, A.; Ahsen, V.; Gul, A.; Bekaroglu, O. J. Chem. Soc. Dalton, 1991, 3367; Okur, A.I.; Gul, A.; Cihan, A.; Tan, N.; Bekaroglu, O. Synth. React. Inorg. Metal-Org. Chem. 1990, 20, 1399. Kobayashi, N.; Opallo, M.; Osa, T. Heterocycles, 1990, 30, 389. Ahsen, V.; Yilmazer, E.; Ertas, M.; Bekaroglu, O. J. Chem. Soc. Dalton 1988, 401. Ahsen, V.; Yilmazer, E.; Bekaroglu, O.; Gul, A. MAkromol. Chem. 1987 8, 243. Ahsen, V.; Yilmazer, E.; Gul, A.; Bekaroglu, O. J. Chem. Res. (S) 1988 234. Ahsen, V.; Yilmazer, E.; Bekaroglu, O.. Makromol. Chem. 1988 189, 2533. Ahsen, V.; Gurek, A. Musluoglu, E.; Bekaroglu, O.; Kocaeli, T. Chem. Ber. 1989, 122, 1073. Ahsen, V.; Yilmazer, E.; Gurek, A.; Gul, A.; Bekaroglu, O.; Gebze, T. Helv. Chim. Acta 1988, 71, 1616. Anzai, J.I.; Liu, C.C. Polymer Comm 1990, 31, 223. Bardin, M.; Bertounesque, E.; Plichon, V.; Simon, J.; Ahsen, V.; Bekaroglu, O., J. Electroanal. Chem. Interfacial Electrochem. 1989, 271, 173. Sielcken, Ot E.; van Tilborg, M. M.; Roks, M. F. M.; Hendriks, R.; Drenth, W.; Nolte, J. M. J. Am. Chem. Soc. 1987, 109, 4261. Sielcken, Ot E.; Schram, J.; Nolte, R. J. M.; Schoonman, J.; Drenth, W. Chem. Commun. 1988, 108. Sielcken, Ot E.; Van de Kuil, L. A.; Drenth, W.; Nolte, R. J. M. Chem. Commun. 1988, 1232. Sielcken, Ot E.; Drenth, W.; Nolte, R. J. M. Recl. Trav. Chir.

- Pays-Bas 1990, 109, 425. Sielcken, O.E.; Vandekuil, L.A.; Drenth, W.; Schoonman, J.; Nolte, R. J. M. J. Am. Chem. Soc. 1990, 112, 3086. Sielcken, O. E.; Van Lindert, H. C. A.; Drenth, W.; Schoonman, J.; Schram, J.; Nolte, R. J. M., Ber. Bunsen-Ges. Phys. Chem, 1989, 93 702.
 Sielcken, Ot E.; Van de Kuil, L. A.; Drenth, W.; Nolte, R. J. M. J. Chem. Soc., Chem.
 Commun. 1988, 1232. Koray, A. R.; Ahsen, V.; Bekaroglu, O. J. Chem. Soc. Chem.
 Commun. 1986, 932. Sirlin, C.; Bosio, L.; Simon, J.; Ahsen, V.; Yilmazer, E.; Bekaroglu, O.
 Chem. Phys. Lett. 1987, 139, 362.
- Gunter, M. J.; Johnston, M. R.; <u>Tetrahedron Lett.</u> 1990, 31, 4801. Zhilina, Z. I.; Mel'nik, V. I.; Andronati, S. A. <u>Ukr. Khim. Zh.</u> 1989, 55, 1211. Dasgupta, D.; Thanabal, V.; Krishnan, V., <u>Biochem. Biophys. Res. Commun.</u> 1982, 104, 1427. Thanabal, V.; Krishnan, V., <u>Inorg. Chem.</u> 1982, 21, 3606. Thanabal, V.; Krishnan, V., <u>Polyhedron</u> 1983, 2, 505. Thanabal, V.; Krishnan, V., <u>J. Am. Chem. Soc.</u> 1982, 104, 3643. Van Willigen, H.; Chandrashekar, T. K., <u>J. Am. Chem. Soc.</u> 1986, 108, 709. Xia, S.; Xiao, N.; Niu, C.; Huang, S., <u>Wuhan Daxue Xuebao</u>, <u>Ziran Kexueban</u> 1984, 3, 77. Kobayashi, N.; Osa, T. <u>Heterocycles</u> 1981, 15, 675. Richardson, N. M.; Sutherland, I. O.; Camilleri, P.; Page, J. A.; <u>Tetrahedron Lett.</u> 1985, 26, 3739.
- 13. Janda, P.; Kobayashi, N.; Auburn, P. R.; Lam, H.; Leznoff, C. C.; Lever, A. B. P. <u>Can. J. Chem.</u> 1989, 67, 1109.
- 14. Van der Putten, A.; Elzing, A.; Visscher, W.; Barendrecht, E. J. Electroanal. Chem. 1986, 214, 523.
- 15. Wan, G. X.; Shigehara, K.; Tsuchida, E.; Anson, F. C. J. Electroanal. Chem. 1984, 179, 239.
- 16. Behret, H.; Clauberg, W.; Sandstede, G. Ber. Bunsenges. Phys. Chem. 1979, 83, 139.
- 17. Kobayashi, N.; Lam, H.; Nevin, W. A.; Janda, P.; Leznoff, C. C.; Lever, A. B. P. <u>Inorg.Chem.</u> 1990, 29, 3415.
- 18. Bard, A. J.; Faulkner, L. R. "Electrochemical Methods", John Wiley, New York, 1980.
- 19. Zecevic, S.; Simic-Glavaski, B.; Yeager, E.; Lever, A.B.P.; Minor, P.C. <u>J. Electroanal. Chem.</u>
 1985, 196, 339.
- 20. Nevin, W. A.; Hempstead, M. R.; Liu, W.; Leznoff, C. C.; Lever, A. B. P. Inorg. Chem. 1987, 26,

570.

- 21. Durand, R.R.; Anson, F.C. J. Electroanal. Chem. 1982, 134, 273.
- 22. Levich, V.G. 'Physicochemical Hydrodynamics', Prentice-Hall: Englewood Cliff NJ. 1962; Koutecky, J.; Levich, V.C. Zh. Fiz. Khim. 1956, 32, 1565.
- 23. Koryta, J.; Dvorak, J. "Principles of Electrochemistry", John Wiley and Sons, New York, 1987, p.343.
- 24. (a) Oyama, N.; Oki, N.; Ohno, H.; Ohnuki, Y.; Matsuda, H.; Tsuchida, E. <u>J. Phys. Chem.</u> 1983, 87, 3642. (b) Andrieux, C.P.; Saveant, J.M. <u>J. Electroanal. Chem.</u> 1982, 134, 163; 1982, 142, 1.
 (c) Andrieux, C.P.; Dumas-Barchiat, J. M.; Saveant, J.M. <u>J. Electroanal. Chem.</u> 1982, 131, 1.
- 25. Golovin, M. N.; Seymour, P.; Jayaraj, K.; Fu, Y. S.; Lever, A. B. P. Inorg. Chem. 1990, 29, 1719.
- 26. Cotton, F. A.; Wilkinson G. "Modern Inorganic Chemistry", 5th Edition, John Wiley, New York, 1987.
- 27. Cariati, F.; Gallizzioli, D.; Morazzoni, F, Busetto, C, <u>J. Chem. Soc. Dalton</u> 1975, 556. Cariati, F.; Morazzoni, F.; Busetto, C, <u>J. Chem. Soc. Dalton</u>, 1976, 496.
- 28. Ni, C-L.; Anson, F. C. <u>Inorg. Chem.</u> 1985, <u>24</u>, 4754.

Legends

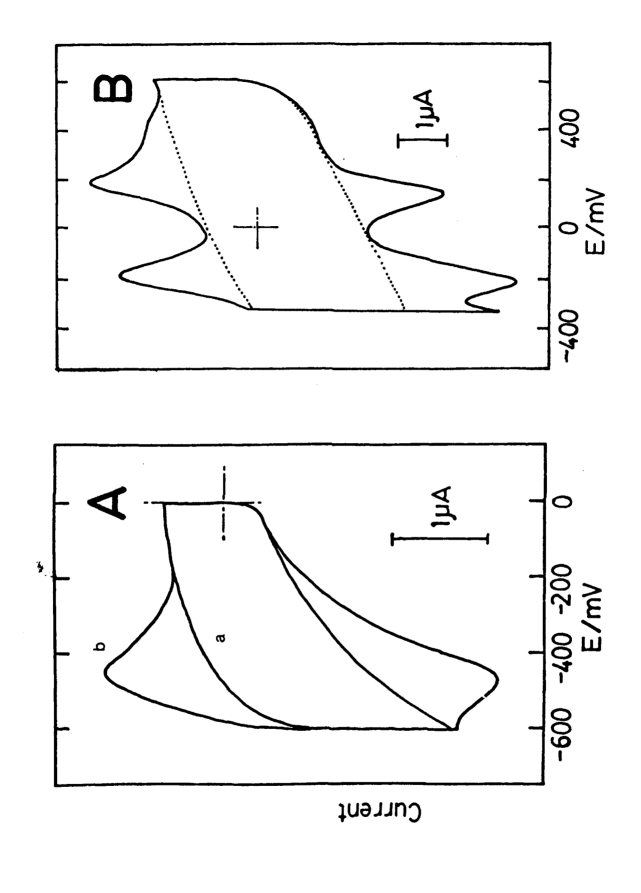
- Figure 1. Cobalt and iron Crown phthalocyanines
- Figure 2. CV Diagrams of (A) Co^{II}CRPc(-2)/HOPG at pH 3.0 and (B) Fe^{II}CRPc(-2)/HOPG at pH 0.9. Dotted lines indicate CV response at a bare HOPG surface. Scan speed 100 mV/s.
- Figure 3. pH Dependencies of (A) oxygen reduction potentials at Co^{II}CRPc(-2)/HOPG, (B)

 Co^{II}CRPc(-2)/[Co^ICRPc(-2)]⁻ couple at HOPG, (C) [Co^ITsPc(-2)]⁻/[Co^ITsPc(-3)]²⁻ couple at an ordinary pyrolytic graphite (OPG) electrode, and (D) hydrogen peroxide reduction at Co^{II}CRPc(-2)/HOPG.
- Figure 4. pH Dependencies of (A) [Fe^{III}CRPc(-2)] +/Fe^{II}CRPc(-2) at HOPG. (B) Oxygen reduction potentials at Fe^{II}CRPc/HOPG from CV peak potentials. (C)

 Fe^{II}CRPc(-2)/[Fe^ICRPc(-2)] couples at HOPG.
- Figure 5. (A) CV curves for oxygen reduction at Co^{II}CRPc/HOPG, scan rate 100mV/s. (B) RDE responses for oxygen reduction at Co^{II}CRPc/HOPG at various pH values, scan rate 10 mV/s. (C) RDE responses for oxygen reduction at Co^{II}CRPc/HOPG at various rotation speeds, pH 13, scan rate 10 mV/s. (D) Koutecky-Levich plots constructed from the plateau current in (D). (E) Tafel type plots from RDE data at Co^{II}CRPc/HOPG for the first oxygen reduction process.
- Figure 6. (A) CV diagrams at Fe^{II}CRPc/HOPG under argon (curve a) and in oxygen saturated solution (curve b) at pH 8.9. Dotted lines indicate responses at bare HOPG, scan rate 100 mV/s. (B) RDE responses at Fe^{II}CRPc/HOPG in oxygen saturated solution at pH 9.2, scan rate 10 mV/s.

M=Co:CoCRPc M=Fe:FeCRPc

Figure 1



igure .

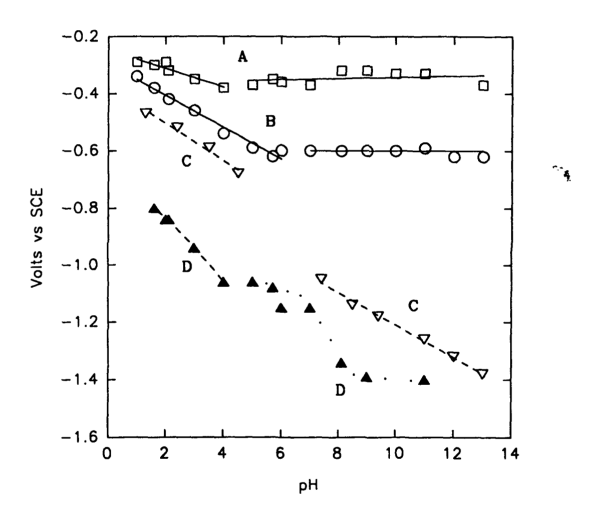


Figure 3

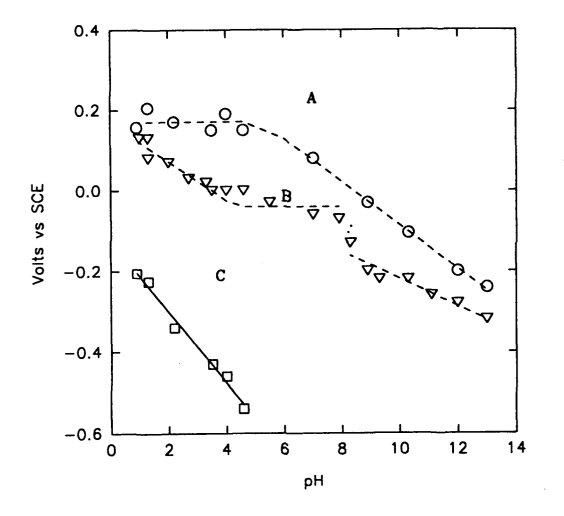


Figure 4

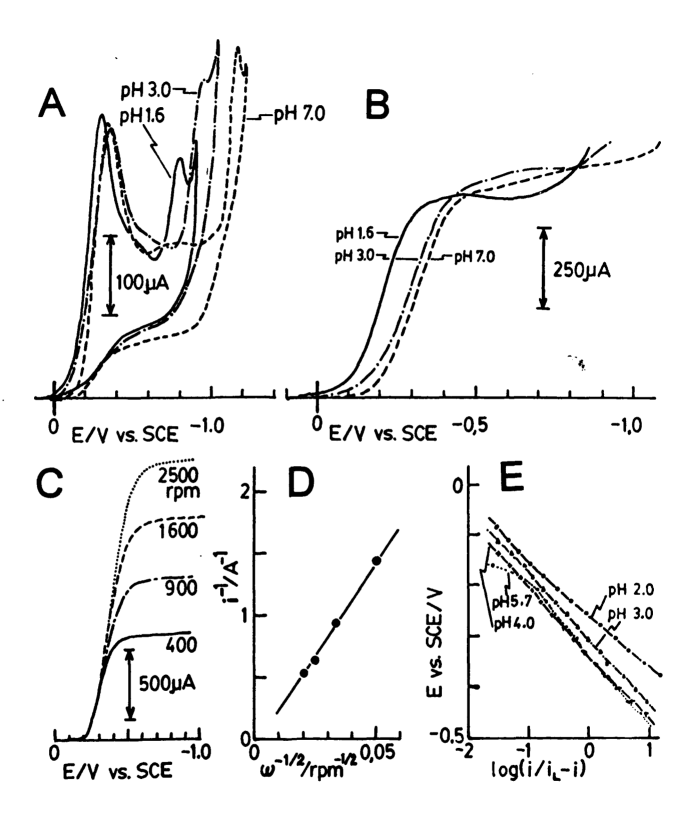


Figure 5

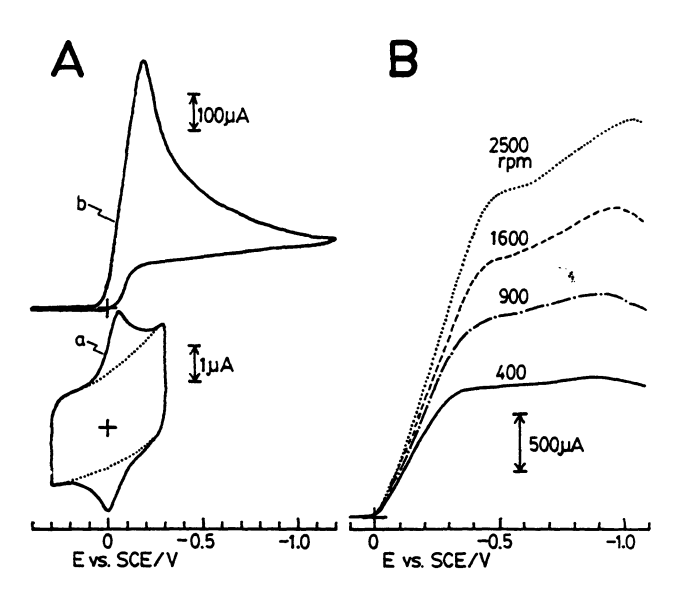


Figure 6